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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

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OFFICE OF
PESTICIDES AND TOXIC SUBSTANCES

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MEMORANDUM

Sostram Chemical Co. Response to the Simazine SUBJECT:

Reregistration Standard: Product Chemistry Data (MRID

#41301901, DEB# 6237.)

FROM: R. B. Perfetti, Ph.D., Chemist

Reregistration Section Dietary Exposure Branch

Health Effects Division (H7509C)

THRU: W. J. Boodee, Section Head

Reregistration Section Dietary Exposure Branch

Health Effects Division (H7509C)

TO: Reto Engler, Ph.D., Chief

Science Analysis and Coordination Branch

Health Effects Division (H7509C)

and

J. Talarico, RM-74 Reregistration Branch

Special Review and Reregistration Division

Attached is the review of product chemistry data submitted by Sostram Chemical Co. in response to the simazine reregistration This information was reviewed by Dynamac Corporation under supervision of Dietary Exposure Branch, HED.

The due date for this review was January 20, 1990.

This information has undergone secondary review in Dietary Exposure Branch and have been revised to reflect the Branch policies.

Please note that Attachment 2 to this review contains Confidential Appendices A, B, C, and , D. These are to be

protected. Copies of these appendices wil be sent only to Toxicology Branch, Reregistration Branch (J. Talarico) and FOD (J. Burrell). All other copies will remain in this Branch.

Please note that revised Tables A and B are attached at the end of the review.

If you need additional input please advise.

Attachment 1: Review of Simazine Product Chemistry Data Attachment 2: Confidential Appendices A,B,C and D

- CC: With Attachments 1 and 2: R.B. Perfetti, R. Coberly (TOX), J. Burrell (FOD), Ethoprop Reregistration Standard File, Ethoprop Subject File, J. Talerico (RB, SR&RD)
- CC: Without Attachments: P. Fenner-Crisp (HED), M. Hawkins (HED), F. Bishop (RD), Circulation (7), RF, R. Engler (SACB)

H7509C:DEB:X77484:CM#2:RM810:R.B.Perfetti:rp:1/24/89



Final Report

SIMAZINE Task 4: Registrant's Response to Product Chemistry Data Requirements

January 24, 1990

Contract No. 68-D8-0080

Submitted to: Environmental Protection Agency Arlington, VA 22202

Submitted by:
Dynamac Corporation
The Dynamac Building
11140 Rockville Pike
Rockville, MD 20852

SIMAZINE

REGISTRANT'S RESPONSE TO PRODUCT CHEMISTRY DATA REQUIREMENTS

Task - 4

BACKGROUND

In response to the Simazine Guidance Document, dated April 14, 1984, Sostram Chemical Company, a subsidiary of Oxon Italia S.P.A. of Italy, has submitted updated product chemistry data (1989; MRID 41301901) for the 95% technical (T) product (EPA Reg. No. 35915-10). These data and our conclusions are discussed below.

61-1. Product Identity and Disclosure of Ingredients

A Confidential Statement of Formula (CSF) has been submitted by the registrant (see Confidential Appendix A). These data do not satisfy the requirements of 40 CFR §158.155 (Guidelines Reference No. 61-1) regarding product composition for the Sostram 95% T (EPA Reg. No. 35915-10) because the CAS registry number and CA-approved chemical name was not provided for the active ingredient and impurities were incorrectly identified as inerts. Additional data are required for this topic.

61-2. Description of Beginning Materials and Manufacturing Process

The registrant has submitted the names and addresses of the suppliers, but not the specifications of the starting materials. The registrant has also submitted a description of the manufacturing process for the technical material (MRID 41301901). This information is reviewed in Confidential Appendix B. These data do not satisfy the requirements of 40 CFR §158.160 (Guideline Reference No. 61-2) regarding starting materials for the Sostram 95% T (EPA Reg. No. 35915-10) because the registrant did not describe the type of process (batch or continuous); furthermore, the registrant did not report specifications of starting materials, the pH and duration of the reaction, the identity of a solvent, quality control steps, or a description of the the process equipment. Additional data are required.

61-3. Discussion of Formation of Impurities

The registrant reported the name of an impurity formed during the synthesis of the technical product (MRID 41301901); this information is presented in Confidential Appendix C. This information does not satisfy the requirements of 40 CFR §158.167 (Guidelines Reference No. 61-3) regarding formation of impurities in the Sostram 95% T (EPA Reg. No. 35915-10) because the formation of the impurity was not adequately described and no

discussions were presented regarding carryover of impurities from the starting materials, solvent, or process intermediates. In addition, there was no discussion of post-production contamination or nitrosamine formation. Additional data are required.

62-1. Preliminary Analysis

The registrant has provided preliminary analyses of five batches each of the registered technical (MRID 41301901). The results of the preliminary analyses are discussed in Confidential Appendix D. These data do not satisfy the requirements of 40 CFR §158.170 (Guideline Reference No. 62-1) regarding preliminary analysis for the Sostram 95% T (EPA Reg. No. 35915-10) because no validation data or statements of precision and accuracy were provided for the methods used to obtain these data. Furthermore, no enforcement analytical method was submitted for we note that no data were reported on the analytical statements of the submitted for

We note that no data were reported on the analysis of the product for nitrosamines. Additional data are required.

62-2. Certification of Limits

Certified limits reported in the CSF are presented in Confidential Appendix A. These data do not satisfy the requirements of 40 CFR §158.175 (Guidelines Reference No. 62-2) regarding certified limits for the Sostram 95% T (EPA Reg. No. 35915-10) because upper and lower limits were transposed between columns on the CSF. Furthermore, an explanation of how the certified limits were established for impurities was not provided. Additional data must be submitted.

62-3. Analytical Methods to Verify Certified Limits

No validation data or statements of precision and accuracy were reported. This description does not satisfy the requirements of 40 CFR §158.180 (Guideline Reference No. 62-3) regarding enforcement analytical methods for the Sostram 95% T (EPA Reg. No. 35915-10) because the registrant did not report validation data or a statement of precision and accuracy of the method.

The registrant has submitted analytical methods for the determination of other impurities in the technical product (MRID 41301901). The procedures are discussed in Confidential Appendix E. This information not satisfy the requirements of 40 CFR §158.180 (Guideline Reference No. 62-3) regarding enforcement

analytical methods for the Sostram 95% T (EPA Reg. No. 35915-10) because the registrant did not submit a description and validation information of a method to determine an impurity of toxicological significance. Additional data are required.

PHYSICAL AND CHEMICAL CHARACTERISTICS

The physical and chemical characteristics for the technical product have been submitted by the registrant (MRID 41301901). These properties are presented in Table 1. Information submitted on color, physical state, odor, melting point, density, solubility, pH, and explodability are adequate. Data on the octanol/water partition coefficient are not adequate because the registrant did not specify whether the test material was pure active ingredient. No data were submitted for vapor pressure, stability, oxidizing or reducing action, storage stability, or corrosiveness. This constitutes a data gap.

Table 1. Physical and chemical properties of the simazine 95% T (EPA Reg. No. 35915-10). All data pertain to the technical grade of the active ingredient (TGAI; MRID 41301901).

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No.,	elines Reference 40 CFR §158.190; of Property	Description [Method]
63-2.	Color	white
63-3.	Physical state	homogeneous powder
63-4.	Odor	odorless
63-5.	Melting point	227-230 C
63-6.	Boiling point	Simazine is a solid at room temperature.
63-7.	Density, bulk density, or specific gravity	450-500 g/l
63-8.	Solubility	SolventSolubilityTemperaturewater5 mg/l20 Cmethanol400 g/kg20 Cchloroform900 g/kg20 C
63-9.	Vapor pressure	not submitted
63-10.	Dissociation constant	data not submitted
53-11.	Octanol/water partition coefficient	$K_{OH} = 143 \text{ (log P = 2.15) [OECD Guideline]}$
3-12.	рН	7.5-9.5
	_	

Table	1.	(Continued.)
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Guidelines Reference No., 40 CFR §158.190;	
Name of Property	Description [Method]
63-13. Stability	data not submitted
63-14. Oxidizing or reducing action-	data not submitted
63-15. Flammability	not required for solids
63-16. Explodability	not explosive
33-17. Storage stability	no data submitted
3-18. Viscosity	simazine T is a solid at room temperature.
3-19. Miscibility	simazine T is a solid at room temperature.
3-20. Corrosiveness	not corrosive

TABLE A. GENERIC DATA REQUIREMENTS FOR THE SIMAZINE TECHNICAL GRADE OF THE ACTIVE INGREDIENT.

1

Data Requirement	Test Substance ²	Guidel ine Status	Must additional data be submitted under FIFRA Sec. 3(c)(2)(B)? [Yes] [No]	Reference (MRID No.)
40 CFR §158.155-190 Product Chemistry Product Composition	mistry	-		
61-2. Beginning Materials & Production Process	TGAL	œ	, **	41301901
61-3. Formation of Impurities	TGAI	æ	*	41301901
Analysis and Certification of Prod 62-1. Preliminary Analysis	roduct Ingredients TGAI	ম মূ	፟፠	41301901
Physical and Chemical Characterist	istics			
63-2. ©lor	TCAI	2	*	1001001
	TCAI	~	×	41301901 41301901
	TGAI	~	: >	41301001
	TGAI	œ	; ≻	41301901 41301001
	TGAI	N/A ⁷	: ×	TOSTOCTA
63-7. Density, Bulk Density, or Specific Gravity		`¤	:×	41301901
	TGAI or PAI	Ω	>	
	ö	€		41301901 N/A
	or	i es	: ×	W/N
63-11. Octanol/Water Partition	PAI	8	: ሜ	41301901
Coefficient				
	TCAI	F	×	41301901
63-13. Stability	TCAI	~	×	N/A
~				
64-1. Submittal of Samples	TGAI or PAI	F	°×	N/A

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- Additional data requirements are listed in the following Table B, "Generic Data Requirements for the Generic data requirements pertain to the TGAI of the Sostram Chemical Co. 95% T (EPA Reg. No. 35915-Simazine 95% T (EPA Reg. No. 35915-10)", a registered technical product.
- Test substance: MP = manufacturing-use product; PAI = purified active ingredient; TEP = typical end-use product; TGAI = technical grade of the active ingredient.
- purification procedures (including procedures to recover or recycle starting materials, intermediates or the process; (iii) a description of the equipment used; (iv) a description of the conditions (e.g., temperature, production process (e.g., batch or continuous); (ii) a description of a solvent used in the manufacturing pressure, pH, humidity) that are controlled during each step of the process; and (v) a description of the (vi) information concerning the composition of each starting material; and, (vii) a description of any procedures used to assure consistent composition of the substance produced (quality control methods); 3. The following information must be provided: (i) a general characterization of the formulation or
- conducted by or for the registrant, and (ii) each impurity which the registrant has reason to believe may be intended and side reactions which may occur during production, the possible degradation of ingredients after impurity associated with the active ingredient which was found to be present in any analysis of the product production, post-production reactions between ingredients, possible contamination from packaging materials present at a level equal to or greater than 0.1% (w/w) based on the composition of each starting material, 4. A discussion regarding the origin of the following potential impurities must be provided: (i) each or production equipment, and process control, purification and quality control measures.
 - The registrant must provide complete and detailed descriptions of the methods used for sample analysis including statements of their precision and accuracy.
- Subdivision D, Guidelines Reference Nos. 63-2 through 63-13, data must be submitted on physicochemical characteristics (color, physical state, odor, melting point, boiling point, specific gravity, solubility, vapor pressure, dissociation constant, octanol/water partition coefficient, pH, and stability). There additional data requirements listed in Table B pertaining to physicochemical characteristics of those 6. As required by 40 CFR §158.190 and more fully described in the Pesticide Assessment Guidelines, technical products which are also manufacturing use products.
- Data on boiling point are not required since the technical product is a solid at room temperature.
 - The registrant must state whether the data reflect tests on the PAI.
- 9. If samples are required, the Agency will request them.

PRODUCT SPECIFIC DATA REQUIREMENTS FOR SIMAZINE MANUFACTURING-USE PRODUCTS. TABLE B.

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Reference (MRID No.)	41301901	41301901	41301901 41301901 41301901	41301901 41301901 41301901 41301901	41301901 N/A N/A 41301901 N/A N/A 41301901	
Must additional data be submitted under FIFRA Sec. 3(c)(2)(B)? [Yes] [No]	**	፠	፞ ፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟፟	***	x x x x x x x x x x x x x x x x x x x	
Guidel ine Status	88 88	æ	its R R R	& & & &	555××55	
Test Substance ²	Stry MP MP	ΨĐ	duct <u>Ingredien</u> MP MP	SSI FIFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	로 한 한 한 한 한 한 한	
Data Requirement	40 CFR §158.155-190 Product Chemistry Product Composition 61-1. Product Composition 61-2. Beginning Materials & Production/Formulation	3. Formation of Impurities	Analysis and Certification of Product Ingredients 62-1. Preliminary Analysis MP 62-2. Certified Limits MP 62-3. Enforcement Method MP	Physical and Chemical Characteristics 63-2. Color 63-3. Physical State 63-4. Odor 63-7. Density, Bulk Density, or Specific Gravity		
Dat	40 CF Produx 61-1. 61-2.	61-3.	Analys 62-1. 62-2. 62-3.	Physi 63-2. 63-3. 63-7.	63-12. 62-14. 62-15. 63-16. 63-17. 63-19.	

(Continued, footnotes follow)

TABLE B. (Continued).

Reference (MRID No.)	<u> </u>
Must additional data be submitted under FIFRA Sec. 3(c)(2)(B)? [Yes] [No]	7
Guideline Status	g
Test Substance	Ψ
Data Requirement	Other Requirements: 64-1. Submittal of Samples

Additional data requirements are listed in the preceding Table A, "Generic Data Requirements for the Simazine Technical Grade of the Active Data pertain to the Sostram 95% T (EPA Reg. No. 35915-10). Ingredient", for the TGAI of the same product.

Test substance: MP = manufacturing-use product; PAI = purified active ingredient; TEP = typical end-use product; TGAI = technical grade of the active ingredient.

ingredient; furthermore, the registrant must amend the CSF so that all impurities are identified as such. The registrant must report the CAS registry number and CA-approved chemical name of the active

purification procedures (including procedures to recover or recycle starting materials, intermediates or the process; (iii) a description of the equipment used; (iv) a description of the conditions (e.g., temperature, pressure, pH, humidity) that are controlled during each step of the process; and (v) a description of the production process (e.g., batch or continuous); (ii) a description of a solvent used in the manufacturing (vi) information concerning the composition of each starting material; and, (vii) a description of any procedures used to assure consistent composition of the substance produced (quality control methods); (i) a general characterization of the formulation or The following information must be provided: substance produced).

conducted by or for the registrant, and (ii) each impurity which the registrant has reason to believe may be impurity associated with the active ingredient which was found to be present in any analysis of the product possible degradation of ingredients in the product after production, post-production reactions between the ingredients in the product, possible contamination from packaging materials or production equipment, and 5. A discussion regarding the origin of the following potential impurities must be provided: (i) each present in the product at a level equal to or greater than 0.1% (w/w) based on the composition of each starting material, intended and side reactions which may occur in the production of the product, the process control, purification and quality control measures.

- including statements of their precision and accuracy. In addition, all nitrosamines must be identified and quantified by methods sensitive to 1 ppm of N-nitroso contaminants in six samples of each manufacturing-use The registrant must provide complete and detailed descriptions of the methods used for sample analysis product; two samples of each must be analyzed shortly after production, two at 3 months after production, and two at 6 months after production. Upper limits must be proposed for all nitrosamines found.
 - quantitative estimate based on the amounts of ingredients used, etc.) along with information on the accuracy production.) In addition, the CSF must be amended so that upper and lower limits are no longer transposed The registrant must provide an explanation of how the certified limits reported on the Confidential and precision of any analytical procedures used. (Certified limits should be based on the sources and Statement of Formula (CSF) were established (sample analysis using a validated analytical procedure, magnitude of variability in the manufacturing process and the stability of the ingredients following
- quantitative estimate based on the amounts of ingredients used, etc.) along with information on the accuracy production.) In addition, the CSF must be amended so that upper and lower limits are no longer transposed The registrant must provide an explanation of how the certified limits reported on the Confidential and precision of any analytical procedures used. (Certified limits should be based on the sources and Statement of Formula (CSF) were established (sample analysis using a validated analytical procedure, magnitude of variability in the manufacturing process and the stability of the ingredients following
- oxidizing or reducing action, flammability, explodability, storage stability, viscosity, miscibility, and manufacturing-use products which contain only the technical grade of the active ingredient are listed in Subdivision D, Guidelines Reference Nos. 63-2 through 63-20, data must be submitted on physicochemical characteristics of each manufacturing-use product (color, physical state, odor, specific gravity, pH, As required in 40 CFR §158.190 and more fully described in the Pesticide Assessment Guidelines, corrosion characteristics). Additional data requirements regarding physicochemical properties of Table A, "Generic Data Requirements for the Simazine Technical Grade of the Active Ingredient."
 - Data are required on oxidizing/reducing potential if product contains an oxidizing or reducing agent. 10.
 - Data are not required on flammability since the product does not contain combustible liquids,
 - Data on viscosity are required only if the product is a liquid.
- Data on miscibility are required only if the product is an emulsifiable liquid and is to be diluted petroleum solvents. with
- 14. If samples are required, the Agency will request them.

Page is not included in this copy. Pages through/9 are not included in this copy.
The material not included contains the following type of information:
X Identity of product inert ingredients
X Identity of product impurities
X Description of the product manufacturing process
$\frac{\chi}{}$ Description of product quality control procedures
Identity of the source of product ingredients
Sales or other commercial/financial information
A draft product label
The product confidential statement of formula
Information about a pending registration action
FIFRA registration data
The document is a duplicate of page(s)
The document is not responsive to the request
The information not included is generally considered confidential by product registrants. If you have any questions, please contact the individual who prepared the response to your request.